#### => d his

(FILE 'HOME' ENTERED AT 17:57:21 ON 16 JAN 2004)

FILE 'CAPLUS' ENTERED AT 18:00:09 ON 16 JAN 2004

E SCHLOEMER G/IN

17 S E4-E6

Ll L2 0 S L1 AND ACETAMIDE?

0 S L1 AND IMIDAZO? LЗ

7 S L1 AND PROCESS L4 5 S DIMETHYLGLYOXYLAMIDE L5

SELECT L5 5 RN

FILE 'REGISTRY' ENTERED AT 18:16:26 ON 16 JAN 2004 L6 3 S E1-E3

FILE 'CAPLUS' ENTERED AT 18:18:30 ON 16 JAN 2004 SELECT L5 4 RN

FILE 'REGISTRY' ENTERED AT 18:18:53 ON 16 JAN 2004 L7 3 S E4-E6

FILE 'REGISTRY' ENTERED AT 18:30:24 ON 16 JAN 2004  $r_8$ STRUCTURE UPLOADED

L9 0 S L8

FILE 'BEILSTEIN' ENTERED AT 18:31:02 ON 16 JAN 2004

L10 0 S L8 L11 1 S L8 SSS FULL

FILE 'REGISTRY' ENTERED AT 18:32:54 ON 16 JAN 2004 2 S L8 SSS FULL

FILE 'CAPLUS' ENTERED AT 18:33:36 ON 16 JAN 2004 L13 4 S L12

=> d 18

L8 HAS NO ANSWERS

STR OH

G1 Me, Et, n-Pr, i-Pr, n-Bu, i-Bu, s-Bu, t-Bu

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=> d 1-4 bib abs hitstr
      ANSWER 1 OF 4 CAPLUS COPYRIGHT 2004 ACS on STN
      1991:101100 CAPLUS
DN
      One-pot synthesis of N,N,N',N'-tetrasubstituted ureas and oxomalonamides
ΤI
      by oxidative carbonylation of lithium amides at atmospheric pressure
ΑIJ
      Nudelman, Norma S.; Lewkowicz, Elizabeth S.; Perez, Daniel G.
      Fac. Cienc. Exactas, Univ. Buenos Aires, Buenos Aires, 1428, Argent.
CS
      Synthesis (1990), (10), 917-20
CODEN: SYNTBF, ISSN: 0039-7881
SO
DΤ
      Journal
      English
LΑ
      CASREACT 114:101100
OS
      N,N,N',N'-tetrasubstituted ureas RR1NCONRR1 (R = R1 = Bu, cyclohexyl,
AB
      CHMe2, cyclohexyl) were prepd. in good yields by reaction of lithium
      aliph. amides RR1NLi in THF soln. with CO under mild conditions
      (0.degree , 1013 mbar) followed by treatment with oxygen prior to work up.
      N,N,N',N'-tetrasubstituted oxomalonamides (oxopropanediamides)
      RRINCOCOCONRRI were prepd. under similar reaction conditions by carrying
      out the reaction in the presence of known amts. of the pure amine.
      Besides being an useful synthetic method, the present studies afford new
      evidence of the mechanism of the reaction.
TΤ
      83862-73-1P
      RL: SPN (Synthetic preparation); PREP (Preparation)
         (prepn. of)
RN
      83862-73-1 CAPLUS
     Acetamide, 2,2'-oxybis[N,N-dibutyl-2-hydroxy- (9CI) (CA INDEX NAME)
           O OH
                     OH O
(n-Bu) 2N-C-CH-O-CH-C-N (Bu-n) 2
L13
     ANSWER 2 OF 4 CAPLUS COPYRIGHT 2004 ACS on STN
      1988:55481 CAPLUS
     Carbon-carbon bond formation through the carbonylation of lithium
      dialkylamides. One-pot synthesis of N-alkyl-substituted formamides,
      glyoxylamides, and hydroxymalonamides
      Perez, Daniel G.; Nudelman, N. Sbarbati
     Fac. Cienc. Exactas, Univ. Buenos Aires, Buenos Aires, 1428, Argent.
cs
     Journal of Organic Chemistry (1988), 53(2), 408-13
      CODEN: JOCEAH; ISSN: 0022-3263
DT
     Journal
      English
LA
os
     CASREACT 108:55481
     The reaction of RRINLi (R = Rl = Bu, pentyl, cyclohexyl; R = iso-Pr, Rl = cyclohexyl; RR1 = 3-oxapentamethylene) with CO to yield RRINCHO, (RRINCOCHOH) 2O, and RRINCOCH (OH) CONRRI (R, Rl = same as above) was examd.
     under a no. of different conditions. Evidence supporting a lithium carbamoyl intermediate for the latter compds. is presented. A general
     procedure for the prepn. of tetraalkylureas, tetraalkyloxalamides, and
     tetraalkyloxomalonamides is given.
IT
     83862-73-1P
     RL: SPN (Synthetic preparation); PREP (Preparation)
     (prepn. of)
83862-73-1 CAPLUS
RN
CM
     Acetamide, 2,2'-oxybis[N,N-dibutyl-2-hydroxy- (9CI) (CA INDEX NAME)
                     он о
(n-Bu) 2N-C-CH-O-CH-C-N(Bu-n) 2
L13 ANSWER 3 OF 4 CAPLUS COPYRIGHT 2004 ACS on STN
     1983:34210 CAPLUS
AN
DN
     98:34210
     Insertion of carbon monoxide into lithium-nitrogen bonds. One-pot
     synthesis of dialkylformamides and dialkylgloxylamides
     Nudelman, N. Sbarbati; Perez, Daniel
     Fac. Cienc. Exactas Nat., Univ. Buenos Aires, Buenos Aires, 1428, Argent. Journal of Organic Chemistry (1983), 48(1), 133-4
```

CODEN: JOCEAH; ISSN: 0022-3263

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10/620209
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DT
     Journal
     English
LA
     CASREACT 98:34210
os
     Lithium dialkylamides react with CO to afford dialkylformamides (1),
AB
     tetralkylhydroxymalonamides and dialkylglyoxylamides (II). Reaction
     conditions are described to produce I or II in good yields.
IT
     83862-73-1P
     RL: SPN (Synthetic preparation); PREP (Preparation)
        (prepn. of)
     83862-73-1 CAPLUS
RN
CN
     Acetamide, 2,2'-oxybis[N,N-dibutyl-2-hydroxy- (9CI) (CA INDEX NAME)
```

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10/620209
=> s dimethylglyoxylamide
              5 DIMETHYLGLYOXYLAMIDE
=> d 1-5 bib abs kwic
L5
     ANSWER 1 OF 5 CAPLUS COPYRIGHT 2004 ACS on STN
     2003:887680 CAPLUS
AN
DN
     139:364844
     Preparation of indolizines as sPLA2 inhibitors
TI
     Dillard, Robert D.; Hagishita, Sanji; Ohtani, Mitsuaki
IN
PA
     Eli Lilly and Company, USA; Shiongi and Company, Ltd.
SO
     U.S., 62 pp., Cont.-in-part of U.S. Ser. No. 278,445.
     CODEN: USXXAM
תת
     Patent
T.A
     English
FAN. CNT 2
     PATENT NO.
                        KIND DATE
                                                APPLICATION NO. DATE
ΡI
     US 6645976
                         В1
                               20031111
                                                US 1997-765566
     WO 9603383
                              19960208
                                                WO 1995-US9381
                         A1
                                                                  19950720
         W: AM, AT, AU, BB, BG, BR, BY, CA, CH, CN, CZ, DE, DK, EE, ES, FI,
              GB, GE, HU, IS, JP, KE, KG, KP, KR, KZ, LK, LR, LT, LU, LV, MD,
              MG, MN, MW, MX, NO, NZ, PL, PT, RO, RU, SD, SB, SG, SI, SK, TJ,
              TM, TT
          RW: KE, MW, SD, SZ, UG, AT, BE, CH, DE, DK, ES, FR, GB, GR, IE, IT,
              LU, MC, NL, PT, SE, BF, BJ, CF, CG, CI, CM, GA, GN, ML, MR, NE,
              SN, TD, TG
PRAI US 1994-278445
                         A2
                               19940721
     WO 1995-US9381
                               19950720
     MARPAT 139:364844
* STRUCTURE DIAGRAM TOO LARGE FOR DISPLAY - AVAILABLE VIA OFFLINE PRINT *
    Title compds. I, II, III [wherein X = O or S; R11 = independently H,
     alkyl, or halo; R12 = H, halo, (cyclo)alkyl, cycloalkenyl, alkoxy, alkylthio, or a non-interfering substituent having 1-3 atoms other than H;
     R13 = (un) substituted alkyl, alkenyl, alkynyl, (hetero) cyclyl optionally connected by a linking group; R15 and R16 = independently H,
     non-interfering substituent, or (un) substituted (hetero) cyclyl; R17 and
     R18 = independently H, non-interfering substituent, or acidic linker; with
     the proviso that at least one of R17 and R18 must be an acidic linker; or
     pharmaceutically acceptable salt, ester, or amide prodrug derivs.
     thereof], and their 3-acetamide, 3-acetic acid hydrazide, and 3-glyoxylamide analogs were prepd. as inhibitors of human secreted
     phospholipase A2 (sPLA2) mediated release of fatty acids. For example
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conversion of 2-methyl-5-methoxypyridine to the anion in THF using lithium diisopropylamide and subsequent reaction with benzonitrile produced 5-methoxy-2-phenacylpyridine (57.0%). Cyclization of the pyridine deriv. with 1-bromo-2-butanone using NaHCO3 in acetone gave the 1-benzoylindolizine (90.7%), which was reduced by LAH to give 1-benzyl-2-ethyl-6-methoxyindolizine (94.5%). Acylation (98.5%) with Et oxalyl chloride in benzene, followed by sapon. with LiOH in H2O and amidation using NH4OH, provided 2-(1-benzyl-2-ethyl-6-methoxyindolizin-3yl)glyoxylamide. Demethylation by BBr3 in CH2Cl2, coupling with Et 4-bromobutyrate (56.2%) in the presence of NaH in DMF, and hydrolysis with LiOH gave the title indolizine IV (49.9%). Eighty-eight compds. of the invention inhibited recombinant human sPLA2 in a chromogenic assay with IC50 values ranging from 0.006 .mu.M to 1.1 .mu.M, in contrast to IC50 values >50 .mu.M for comparative examples. Administration of 10/mg/kg of the representative compd., 2-[8-(carbomethoxymethoxy)-2-ethyl-3-(2phenylbenzyl) indolizin-1-yl]glyoxylamide, improved the survival rate of male Wistar rats with sPLA2-induced pancreatitis from 33.3% (vehicle) to 91.7%. Thus, invention compds. and their pharmaceutical formulations are useful for the treatment of conditions such as septic shock, adult respiratory distress syndrome, pancreatitis, trauma, bronchial asthma. allergic rhinitis, and rheumatoid arthritis. RE.CNT 6 THERE ARE 6 CITED REFERENCES AVAILABLE FOR THIS RECORD ALL CITATIONS AVAILABLE IN THE RE FORMAT 177556-77-3P, 2-(3-Benzyl-8-hydroxy-2-ethylindolizin-1-yl)acetamide 177556-79-5P, 2-{3-benzyl-5-hydroxy-2-ethylindolizin-1-ylacetamide 177556-79-5P, 2-{2-Ethyl-8-hydroxy-3-(o-phenylbenzyl)indolizin-1-ylacetamide 177556-80-8P, 2-{3-(m-Chlorobenzyl)-2-ethyl-8-hydroxyindolizin-1-ylacetamide 177556-81-9P, 2-{2-Cyclopropyl-8-hydroxy-3-(o-phenylbenzyl)indolizin-1-ylacetamide 177556-84-2P, 2-[8-[[(Benzyloxycarbonyl)methyl]oxy]-2-ethyl-3-(o-phenylbenzyl)indolizin-

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177556-85-3P, 2-[8-[[(Benzyloxycarbonyl)methyl]oxy]-3-(m-
 177556-93-3P, 2-(3-Benzyl-8-benzyloxy-2-ethylindolizin-1-yl)glyoxylamide
177556-94-4P, 2-(3-Benzyl-8-benzyloxy-2-ethylindolizin-1-yl)glyoxylamide
177556-94-4P, 2-(3-Benzyl-8-benzyloxy-2-ethylindolizin-1-yl)glyoxylamide
177556-94-4P, 2-(3-Benzyl-8-benzyloxy-2-ethylindolizin-1-yl)glyoxylamide
 yl)-N-methylqlyoxylamide
                                                                             177556-95-5P, 2-(3-Benzyl-8-benzyloxy-2-
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  2-[8-Benzyloxy-2-ethyl-3-(o-phenylbenzyl)indolizin-1-yl]glyoxylamide
2-[8-Benzyloxy-2-ethyl-3-(o-phenylbenzyl)indolizin-1-yl]glyoxylamide
177556-97-7P, 2-[8-Benzyloxy-2-ethyl-3-(o-phenylbenzyl)indolizin-1-yl]-N-
methylglyoxylamide 177556-98-8P, 2-[8-Benzyloxy-2-ethyl-3-(o-
phenylbenzyl)indolizin-1-yl]-N,N-dimethylglyoxylamide
177556-99-9P, 2-(3-Benzyl-8-benzyloxy-2-methylindolizin-1-yl)glyoxylamide
177557-00-5P, 2-[8-Benzyloxy-3-(m-chlorobenzyl)-2-ethyl-3-(m-
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 phenylbenzyl)indolizin-1-yl]glyoxylamide 177557-26-5P,
2-(8-Benzyloxy-3-cyclohexylmethyl-2-ethylindolizin-1-yl)glyoxylamide
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 ethylindolizin-1-yl)glyoxylamide 177557-29-8P, 2-[8-Benzyloxy-2-ethyl-3-pentylindolizin-1-yl)glyoxylamide 177557-30-1P, 2-[8-Benzyloxy-2-ethyl-3
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  (2-propylpentyl)indolizin-1-yl]glyoxylamide 177557-31-2P,
 2-[8-Benzyloxy-2-ethyl-3-[(naphth-2-yl)methyl]indolizin-1-yl]glyoxylamide 177557-32-3P, 2-[8-Benzyloxy-3-(3,5-di-tert-butylbenzyl)-2-ethylindolizin-
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  2-[8-Benzyloxy-3-(o-benzylbenzyl)-2-ethylindolizin-1-yl]qlyoxylamide
 177557-35-6P, 2-[8-Benzyloxy-2-ethyl-3-[(thiophen-2-yl)methyl]indolizin-1-
 yl]glyoxylamide 177557-36-7P, 2-[8-Benzyloxy-2-ethyl-3-[[3-(thiophen-2-yl)thiophen-2-yl]methyl]indolizin-1-yl]glyoxylamide 177557-37-8P,
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 177557-38-9P, 2-[8-Benzyloxy-2-ethyl-3-(o-nitrobenzyl)indolizin-1-
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177557-51-6P,
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  [[(carbomethoxy)methyl]oxy]-2-methylindolizin-1-yl]glyoxylamide
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cyclopropyl-3-[(1-naphthyl)methyl]indolizin-1-yl]glyoxylamide
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177560-00-8P, 2-[8-[[(Methoxycarbonyl)methyl]amino]-3-cyclohexylmethyl-2-
methylindolizin-1-yl]glyoxylamide 182115-63-5P, Methyl 2-[[3-Benzyl-1-(carbamoylmethyl)-2-ethylindolizin-8-yl]oxy]acetate
182115-76-0P, 2-[8-Benzyloxy-2-ethyl-3-[(4-pentylcyclohexyl)methyl]indoliz in-1-yl]glyoxylamide 182115-78-2P, 2-(8-Benzyloxy-3-cyclopentylmethyl-2-cyclopropylindolizin-1-yl)glyoxylamide 182115-84-0P, 2-[8-Hydroxy-2-ethyl-3-(4-pentylcyclohexylmethyl)indolizin-1-
2-16-hydroxylamide 182115-86-2P, 2-[3-Benzyl-8-[[(carbethoxy)methyl]oxy]-2-ethylindolizin-1-yl]glyoxylamide 182115-87-3P, 2-[8-
[[(Carbethoxy)methyl]oxy]-2-ethyl-3-(o-phenylbenzyl)indolizin-1-
yl]glyoxylamide 182115-88-4P, 2-[8-[[(Carbethoxy)methyl]oxy]-3-(m-
yllglyoxylamide
chlorobenzyl) -2-ethylindolizin-1-yl]glyoxylamide
                                                                 182115-90-8P,
2-[8-[[(Carbethoxy)methyl]oxy]-2-ethyl-3-[(1-naphthyl)methyl]indolizin-1-
yllglyoxylamide 182115-92-0P, 2-[3-Benzyl-8-[[(carbethoxy)methyl]oxy]-2-methylindolizin-1-yllglyoxylamide 182115-93-1P, 2-[8-[[(Carbomethoxy)methyl]oxy]-2-ethyl-3-[(4-pentylcyclohexyl)methyl]indolizi
n-l-yllglyoxylamide
                           182116-42-3P, 2-[7-(5-Carboethoxypentyloxy)-2-ethyl-
3-(o-phenylbenzyl)indolizin-1-yl)glyoxylamide 182116-44-5P,
2-[3-Benzyl-8-[[(methoxycarbonyl)methyl]amino]-2-methylindolizin-1-
yl]acetamide
                  182116-45-6P, 2-[3-Benzyl-8-[(carboxymethyl)amino]-2-
methylindolizin-1-yl]acetamide 182116-49-0P, 2-[8-(3-Carbomethoxypropyloxy)-2-ethyl-3-(o-phenylbenzyl)indolizin-1-
yllglyoxylamide
                      215160-62-6P, 2-[8-[[(Carbomethoxy)methyl]oxy]-2-ethyl-3-
(o-phenylbenzyl)indolizin-1-yl]glyoxylamide 215160-63-7P,
2-[3-Benzyl-8-[[(tert-butoxycarbonyl)methyl]oxy]-2-ethylindolizin-1-
yllglyoxylamide 215160-64-8P 215160-65-9P 622835-99-8P,
2-[3-(1-Naphthyl)-8-hydroxy-2-ethylindolizin-1-yl]acetamide
622836-00-4P, Methyl 2-[[3-Naphthyl-1-(carbamoylmethyl)-2-ethylindolizin-8-
ylloxylacetate 622836-03-7P, 2-[3-Benzyl-8-[[(carbethoxy)methyl]oxy]-2-ethylindolizin-1-yll-N-methylglyoxylamide 622836-04-8P,
2-[3-Benzyl-8-[((carbethoxy)methyl)oxy]-2-ethylindolizin-1-yl]-N,N-dimethylglyoxylamide 622836-05-9P, 2-[8-
[[(Carbethoxy)methyl]oxy]-2-ethyl-3-(o-phenylbenzyl)indolizin-1-yl]-N,N-
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dimethylglyoxylamide 622836-07-1P, 2-{8-(Cyanomethyloxy)-2-ethyl-3-{(1-naphthyl)methyl}indolizin-1-yl}glyoxylamide 622836-33-3P, 2-[3-{(Adamant-1-yl)methyl}-8-benzyloxy-2-ethylindolizin-1-yl}glyoxylamide 622836-34-4P, 8-Benzyloxy-3-(cyclopentylcarbonyl)-2-cyclopropylindolizine
 622836-35-5P, 8-Benzyloxy-3-cyclopentylmethyl-2-cyclopropylindolizine
 622836-36-6P, 2-[8-[[(Carbomethoxy)methyl]oxy]-2-ethyl-3-(thiophen-2-
yl)indolizin-1-yl]glyoxylamide 622836-37-7P, 2-(3-Cyclopentylmethyl-2-
 cyclopropyl-8-hydroxyindolizin-1-yl)glyoxylamide
                                                                                                                   622836~57-1P,
 2-[3-(Biphenyl 2 yl)-8-[[(carbomethoxy)methyl]oxy]-2-methoxyindolizin-1-
yl]glyoxylamide
 RL: PAC (Pharmacological activity); RCT (Reactant); SPN (Synthetic
preparation); THU (Therapeutic use); BIOL (Biological study); PREP
 (Preparation); RACT (Reactant or reagent); USES (Uses)
       (sPLA2 inhibitor; prepm. of indolizines as inhibitors of sPLA2 mediated release of fatty acids)
177556-76-2P, 2-[1-Benzyl-6-(3-carboxypropyloxy)-2-ethylindolizin-3-yl]glyoxylamide 177556-87-5P, 2-[3-Benzyl-8-[(carboxymethyl)oxy]-2-ethylindolizin-1-yl]acetamide 177556-89-7P, 2-[8-[(Carboxymethyl)oxy]-2-
ethylindolizin-1-yl]acetamide 177556-89-7P, 2-[8-[(Carboxymeth ethyl-3-(o-phenylbenzyl)indolizin-1-yl]acetamide 177556-90-0P,
2-[8-[(Carboxymethyl)oxy]-3-(m-chlorobenzyl)-2-ethylindolizin-1-yl]acetamide 177556-91-1P, 2-[8-[(Carboxymethyl)oxy]-2-cyclopropyl-3-(o-
phenylbenzyl) indolizin-1-yl]acetamide 177557-67-4P, 2-[2-Ethyl-8-
[(carboxymethyl)oxy]-3-(p-phenylbenzyl) indolizin-1-yl]glyoxylamide
t(Carboxymethyl) (3-1) - 1-1 - 1-1 - 1-1 - 1-1 - 1-1 - 1-1 - 1-1 - 1-1 - 1-1 - 1-1 - 1-1 - 1-1 - 1-1 - 1-1 - 1-1 - 1-1 - 1-1 - 1-1 - 1-1 - 1-1 - 1-1 - 1-1 - 1-1 - 1-1 - 1-1 - 1-1 - 1-1 - 1-1 - 1-1 - 1-1 - 1-1 - 1-1 - 1-1 - 1-1 - 1-1 - 1-1 - 1-1 - 1-1 - 1-1 - 1-1 - 1-1 - 1-1 - 1-1 - 1-1 - 1-1 - 1-1 - 1-1 - 1-1 - 1-1 - 1-1 - 1-1 - 1-1 - 1-1 - 1-1 - 1-1 - 1-1 - 1-1 - 1-1 - 1-1 - 1-1 - 1-1 - 1-1 - 1-1 - 1-1 - 1-1 - 1-1 - 1-1 - 1-1 - 1-1 - 1-1 - 1-1 - 1-1 - 1-1 - 1-1 - 1-1 - 1-1 - 1-1 - 1-1 - 1-1 - 1-1 - 1-1 - 1-1 - 1-1 - 1-1 - 1-1 - 1-1 - 1-1 - 1-1 - 1-1 - 1-1 - 1-1 - 1-1 - 1-1 - 1-1 - 1-1 - 1-1 - 1-1 - 1-1 - 1-1 - 1-1 - 1-1 - 1-1 - 1-1 - 1-1 - 1-1 - 1-1 - 1-1 - 1-1 - 1-1 - 1-1 - 1-1 - 1-1 - 1-1 - 1-1 - 1-1 - 1-1 - 1-1 - 1-1 - 1-1 - 1-1 - 1-1 - 1-1 - 1-1 - 1-1 - 1-1 - 1-1 - 1-1 - 1-1 - 1-1 - 1-1 - 1-1 - 1-1 - 1-1 - 1-1 - 1-1 - 1-1 - 1-1 - 1-1 - 1-1 - 1-1 - 1-1 - 1-1 - 1-1 - 1-1 - 1-1 - 1-1 - 1-1 - 1-1 - 1-1 - 1-1 - 1-1 - 1-1 - 1-1 - 1-1 - 1-1 - 1-1 - 1-1 - 1-1 - 1-1 - 1-1 - 1-1 - 1-1 - 1-1 - 1-1 - 1-1 - 1-1 - 1-1 - 1-1 - 1-1 - 1-1 - 1-1 - 1-1 - 1-1 - 1-1 - 1-1 - 1-1 - 1-1 - 1-1 - 1-1 - 1-1 - 1-1 - 1-1 - 1-1 - 1-1 - 1-1 - 1-1 - 1-1 - 1-1 - 1-1 - 1-1 - 1-1 - 1-1 - 1-1 - 1-1 - 1-1 - 1-1 - 1-1 - 1-1 - 1-1 - 1-1 - 1-1 - 1-1 - 1-1 - 1-1 - 1-1 - 1-1 - 1-1 - 1-1 - 1-1 - 1-1 - 1-1 - 1-1 - 1-1 - 1-1 - 1-1 - 1-1 - 1-1 - 1-1 - 1-1 - 1-1 - 1-1 - 1-1 - 1-1 - 1-1 - 1-1 - 1-1 - 1-1 - 1-1 - 1-1 - 1-1 - 1-1 - 1-1 - 1-1 - 1-1 - 1-1 - 1-1 - 1-1 - 1-1 - 1-1 - 1-1 - 1-1 - 1-1 - 1-1 - 1-1 - 1-1 - 1-1 - 1-1 - 1-1 - 1-1 - 1-1 - 1-1 - 1-1 - 1-1 - 1-1 - 1-1 - 1-1 - 1-1 - 1-1 - 1-1 - 1-1 - 1-1 - 1-1 - 1-1 - 1-1 - 1-1 - 1-1 - 1-1 - 1-1 - 1-1 - 1-1 - 1-1 - 1-1 - 1-1 - 1-1 - 1-1 - 1-1 - 1-1 - 1-1 - 1-1 - 1-1 - 1-1 - 1-1 - 1-1 - 1-1 - 1-1 - 1-1 - 1-1 - 1-1 - 1-1 - 1-1 - 1-1 - 1-1 - 1-1 - 1-1 - 1-1 - 1-1 - 1-1 - 1-1 - 1-1 - 1-1 - 1-1 - 1-1 - 1-1 - 1-1 - 1-1 - 1-1 - 1-1 - 1-1 - 1-1 - 1-1 - 1-1 - 1-1 - 1-1 - 1-1 - 1-1 - 1-1 - 1-1 - 1-1 - 1-1 - 1-1 - 1-1 - 1-1 - 1-1 - 1-1 - 1-1 - 1-1 - 1-1 - 1-1 - 1-1 - 1-1 - 1-1 - 1-1 - 1-1 - 1-1 - 1-1 - 1-1 - 1-1 -
3-cyclopentylmethyl-2-ethylindolizin-1-yl]glyoxylamide 177557-70-9P,
2-[8-[(Carboxymethyl)oxy]-3-cycloheptylmethyl-2-ethylindolizin-1-
yl]glyoxylamide 177557-71-0P, 2-[8-[(Carboxymethyl)oxy]-2-ethyl-3-
pentylindolizin-1-yl]glyoxylamide 177557-72-1P, 2-[8-
[(Carboxymethyl)oxy]-2-ethyl-3-(2-propylpentyl)indolizin-1-yl]glyoxylamide
177557-75-4P, 2-[8-[(Carboxymethyl)oxy]-2-ethyl-3-(2-phenylethyl)indolizin-
1-yl]glyoxylamide 177557-76-5P, 2-[8-[(Carboxymethyl)oxy]-3-(o-
benzylbenzyl)-2-ethylindolizin-1-yl]glyoxylamide 177557-77-6P,
2-[8-[(Carboxymethyl)oxy]-2-ethyl-3-[(thiophen-2-yl)methyl]indolizin-1-
yl]glyoxylamide 177557-78-7P, 2-[8-[(Carboxymethyl)oxy]-2-ethyl-3-[[3-
                                        177557-78-7P, 2-[8-[(Carboxymethyl)oxy]-2-ethyl-3-[[3-
yl]glyoxylamide
 (thiophen-2-yl)thiophen-2-yl]methyl]indolizin-1-yl]glyoxylamide
177557-79-8P, 2-[2-Ethyl-8-[(carboxymethyl)oxy]-3-(m-
methoxybenzyl)indolizin-1-yl]glyoxylamide 177557-80-1P,
2-[2-Ethyl-8-[(carboxymethyl)oxy]-3-(o-nitrobenzyl)indolizin-1-yl]glyoxylamide 177557-82-3P, 2-[3-[(Adamant-1-yl)methyl]-8-
 [(carboxymethyl)oxy]-2-methylindolizin-1-yl]glyoxylamide
                                                                                                                                       177557-83-4P,
2-[3-Benzyl-8-[(carboxymethyl)oxy]-2-cyclopropylindolizin-1-
yl]glyoxylamide 177557-84-5P, 2-[3-(p-Butylbenzyl)-8-
 [(carboxymethyl)oxy]-2-ethylindolizin-1-yl]glyoxylamide
2-[8-[(Carboxymethyl)oxy]-3-cyclohexylmethyl-2-methylindolizin-1-
yl]glyoxylamide 177557-86-7P, 2-[8-[(Carboxymethyl)oxy]-3-
cyclopentylmethyl-2-cyclopropylindolizin-1-yl]glyoxylamide 177557-87-8P,
 2-[8-[(Carboxymethyl)oxy]-3-cyclopentylmethyl-2-methylindolizin-1-
yl]glyoxylamide 177558-06-4P, 2-[3-[(Biphenyl-2-yl)methyl]-8-
[(carboxymethyl)oxy]-2-ethylindolizin-1-yl]glyoxylamide sodium salt
177558-07-5P, 2-[3-[(Biphenyl-2-yl)methyl]-8-[[[1-(methoxycarbonyloxy)ethoxy]carbonyl]methoxy]-2-ethylindolizin-1-
yl]glyoxylamide
                                       177558-08-6P, 2-[3-[(Biphenyl-2-yl)methyl]-2-ethyl-8-
 [[[1-(isopropyloxycarbonyloxy)ethoxy]carbonyl]methoxy]indolizin-1-
yl]glyoxylamide
                                        177558-11-1P, 2-[3-[(Biphenyl-2-yl)methyl]-8-[[[[[1-
 (cyclopentyloxycarbonyloxy)ethyl]oxy]carbonyl]methyl]oxy]-2-ethylindolizin-
 1-yl]glyoxylamide 177558-12-2P, 2-[3-[(Biphenyl-2-yl)methyl]-8-[([[[1-
((cyclopentylcarbonyl)oxy]ethyl]oxy]carbonyl)methyl]oxy]-2-ethylindolizin-
1-yl]glyoxylamide 177558-18-8P, 2-(3-([Biphenyl-2-yl])methyl]-2-ethyl-8-
[((1H-tetrazol-5-yl)methyl]oxylindolizin-1-yl]glyoxylamide 177558-22-4P
[((1H-tetrazol-5-yl)methyl)oxy]indolizin-1-yl]glyoxylamide 177558-22-4
2-[3-Benzyl-7-(3-carboxypropyloxy)-2-ethylindolizin-1-yl]glyoxylamide
177558-23-5P, 2-[7-(3-Carboxypropyloxy)-2-cyclopropyl-3-(o-phenylbenzyl)indolizin-1-yl]glyoxylamide 177558-24-6P,
2-[7-(3-Carboxypropyloxy)-2-ethyl-3-(o-phenylbenzyl)indolizin-1-yl]glyoxylamide 177558-25-7P, 2-[7-(3-Carboxypropyloxy)-3-cyclohexylmethyl-2-ethylindolizin-1-yl]glyoxylamide 177558-26-8P,
2-[3-[(Biphenyl-2-yl)methyl]-8-(3-carboxypropyloxy)-2-ethylindolizin-1-yl]glyoxylamide 177558-27-9P, 2-[7-[(Carboxymethyl)oxy]-2-ethyl-3-(o-phenylbenzyl)indolizin-1-yl]glyoxylamide 177558-29-1P
 phenylbenzyl)indolizin-1-yllglyoxylamide 177558-29-1P,
2-[3-[(Biphenyl-2-yl)methyl]-8-(2-carboxyethyloxy)-2-ethylindolizin-1-
                                        177558-31-5P, 2-{3-{(Biphenyl-2-yl)methyl}-8-(2-
yl]glyoxylamide
 carbomethoxyethyloxy)-2-ethylindolizin-1-yl]glyoxylamide
2-[7-(3-Carbethoxypropyloxy)-2-ethyl-3-(o-phenylbenzyl)indolizin-1-yl]acetamide 177558-33-7P, 2-[7-(3-Carboxypropyloxy)-2-ethyl-3-(o-phenylbenzyl)-2-ethyl-3-(o-phenylbenzyl)
 phenylbenzyl)indolizin-1-yl]acetamide 177558-35-9P, 2-[8-
 [(Carboxymethyl)oxy]-2-methylthio-3-(o-phenylbenzyl)indolizin-1-
yl]glyoxylamide 177560-01-9P, 2-{3-Benzyl-8-[(carboxymethyl)amino]-2-
 methylindolizin-1-yl]glyoxylamide
                                                                                 177560-02-0P, 2-[8-
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[(Carboxymethyl)amino]-3-cyclohexylmethyl-2-methylindolizin-1-yl]glyoxylamide 182115-96-4P, 2-[3-Benzyl-8-[(carboxymethyl)oxy]-2-ethylindolizin-1-yl]glyoxylamide 182115-97-5P, 2-[8-[(Carboxymethyl)oxy]-
      2-ethyl-3-(o-phenylbenzyl)indolizin-1-yl]glyoxylamide
                                                                         182115-98-6P.
      2-[8-[(Carboxymethyl)oxy]-3-(m-chlorobenzyl)-2-ethylindolizin-1-
     yl|glyoxylamide 182115-99-7P, 2-[8-[(Carboxymethyl)oxy]-2-ethyl-3-(m-
      trifluoromethylbenzyl) indolizin-1-yllglyoxylamide 182116-00-3P.
      2-[8-[(Carboxymethyl)oxy]-2-ethyl-3-[(1-naphthyl)methyl]indolizin-1-
     y||g|yoxylamide 182116-01-4P, 2-[8-[(Carboxymethyl)oxy]-2-cyclopropyl-3-

(o-phenylbenzyl)indolizin-1-yl]glyoxylamide 182116-02-5P,

2-[3-Benzyl-8-[(carboxymethyl)oxy]-2-methylindolizin-1-yl]glyoxylamide
      182116-03-6P, 2-[8-[(Carboxymethyl)oxy]-2-ethyl-3-(4-
     pentylcyclohexylmethyl)indolizin-1-yllglyoxylamide 182116-43-4P.
     2-[7-(5-Carboxypentyloxy)-2-ethyl-3-(o-phenylbenzyl)indolizin-1-yl]glyoxylamide 182116-46-7P, 2-[3-[(Biphenyl-2-yl)methyl]-2-methyl-8-
     [(pyridin-2-yl)methoxy]indolizin-1-yl]glyoxylamide 182116-47-8P, 2-[3-[(Biphenyl-2-yl)methyl]-2-methyl-8-[(pyridin-4-yl)methoxy]indolizin-1-
     yl]glyoxylamide 182116-48-9P, 2-[3-[(Biphenyl-2-yl)methyl]-2-methyl-8-
     [(quinolin-2-y1)methoxy]indolizin-1-y1]glyoxylamide 182116-50-3P, 2-[3-Benzy1-8-[(carboxymethy1)oxy]-2-ethylindolizin-1-y1]-N-
     methylglyoxylamide 182116-51-4P, 2-{3-Benzyl-8-[(carboxymethyl)oxy]-2-ethylindolizin-1-yl}-N.N-dimethylglyoxylamide 622836-01-5P,
      2-[8-[(Carboxymethyl)oxy]-2-ethyl-3-(1-naphthyl)indolizin-1-yl]acetamide
      622836-06-0P, 2-[8-[(Carboxymethyl)oxy]-2-ethyl-3-(o-
      phenylbenzyl) indolizin-1-yl]-N,N-dimethylglyoxylamide
      622836-08-2P, 2-[8-[[(1H-Tetrazol-5-yl)methyl]oxy]-2-ethyl-3-[(1-
     naphthyl)methyl]indolizin-1-yl]glyoxylamide 622836-38-8P,
     2-[8-[(Carboxymethyl)oxy]-2-ethyl-3-(aphth-2-yl)indolizin-1-
yl]glyoxylamide 622836-39-9P, 2-[8-[(Carboxymethyl)oxy]-2-ethyl-3-(4-
     pentylcyclohexylmethyl)indolizin-1-yl]glyoxylamide sodium salt
      622836-40-2P, 2-[8-[(Carboxymethyl)oxy]-3-cyclohexylmethyl-2-
     methylindolizin-1-yl]glyoxylamide sodium salt 622836-41-3P,
     2-[8-[(Carboxymethyl)oxy]-3-cyclopentylmethyl-2-methylindolizin-1-
     yl]glyoxylamide sodium salt 622836-43-5P, 2-[3-[(Biphenyl-2-yl)methyl]-8-
      [[[[(tert-butoxycarbonyl)methyl]oxy]carbonyl]methyl]oxy]-2-ethylindolizin-
      1-yl]glyoxylamide 622836-44-6P, 2-[3-[(Biphenyl-2-yl)methyl]-8-[[[[(1-
      (cyclohexyloxycarbonyl)ethyl]oxy]carbonyl]methyl]oxy]-2-ethylindolizin-1-
     yl]glyoxylamide 622836-45-7P, 2-[3-[(Biphenyl-2-yl)methyl]-2-ethyl-8-
      [[[[1-[(1-methylcyclopentyloxy)carbonyl]ethyl]oxy]carbonyl]methyl]oxy]ind
     olizin-1-yl]qlyoxylamide 622836-46-8P, 2-[3-[(Biphenyl-2-yl)methyl]-2-
     ethyl-8-[[[[[2-(morpholino)ethyl]oxy]carbonyl]methyl]oxy]indolizin-1-
     yl]glyoxylamide 622836-47-9P 622836-48-0P, 2-[3-[(Biphenyl-2-
     y1)methyl]-2-ethyl-8-[[[(2-oxopropyl)oxy]carbonyl]methoxy]indolizin-1-
     yl]glyoxylamide 622836-49-1P, 2-[3-[(Biphenyl-2-yl)methyl]-2-ethyl-8-
[[(1-trityltetrazol-5-yl)methyl]oxy]indolizin-1-yl]glyoxylamide
     622836-50-4P, 2-[7-(2-Carboethoxyethyloxy)-2-ethyl-3-(o-
     phenylbenzyl)indolizin-1-yl)glyoxylamide 622836-58-2P, 2-{3-(Biphenyl-2-yl)-8-{(carboxymethyl)oxy}-2-methoxyindolizin-1-
     yl]glyoxylamide
     RI: PAC (Pharmacological activity); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES
      (Uses)
         (SPLA2 inhibitor; prepn. of indolizines as inhibitors of sPLA2 mediated
         release of fatty acids)
     ANSWER 2 OF 5 CAPLUS COPYRIGHT 2004 ACS on STN
     1988:5780 CAPLUS
     108:5780
      6-(1-carbamoyl-1-hydroxymethyl)penicillanic acid derivatives, their
     preparation, and their use as antibacterial agents and/or .beta.-lactamase
      inhibitors
     Barth, Wayne Ernest
     Pfizer Inc., USA
     Eur. Pat. Appl., 138 pp.
      CODEN: EPXXDW
     Patent
     English
FAN.CNT 1
     PATENT NO.
                          KIND DATE
                                                    APPLICATION NO. DATE
     EP 220939
                                 19870506
                                                    EP 1986-308235
                                                                        19861023
                           Al
          R: AT, BE, CH, DE, ES, FR, GB, GR, IT, LI, LU, NL, SE
     WO 9006928
                                 19900628
                                                    WO 1985-US2134 19851029
                           A1
          W: US
     DK 8605143
                                                    DK 1986-5143
     JP 62142183
                           A2
                                  19870625
                                                    JP 1986-258106
                                                                         19861029
     JP 06092417
                                  19941116
                           B4
     US 4797394
                                  19890110
                                                    US 1987-85675
                                                                         19870605
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US 4868296 19890919 19851029 WO 1985-US2134 PRAI us 1987-85675 19870605 CASREACT 108:5780

os GI

$$R^1R^2NCOCH(OH)$$
 $H$ 
 $S$ 
 $Me$ 
 $Me$ 
 $CO_2R$ 
 $I$ 

III, R3=R4=Br

IV,  $R^3=Br$ ,  $R^4=CH(OH)CO_2CH_2CH=CH_2$ 

US 1988-243568

19880912

V,  $R^3 = H$ ,  $R^4 = CH (OH) CO_2 CH_2 CH = CH_2$ 

VI,  $R^3=H$ ,  $R^4=CH(OH)CO_2H$ 

Title compds. I [n = 0-2; R = H, ester group hydrolyzable under physiol.conditions, acyloxymethyl or 1-(acyloxy)ethyl derived from conventional .beta.-lactam antibiotics; R1, R2 = H, (un)substituted Ph, phenylalkyl, cycloalkyl, naphthyl, azolyl, etc.; NR1R2 = pyrrolidino, piperidino, morpholino, 1,2,3,4-tetrahydroquinolinyl, etc. and their salts, useful as antibacterial agents and/or .beta.-lactamase inhibitors (no data), were prepd. by a) hydrogenolysis of I (R = CH2Ph) and optionally b) converting the compd. to a cationic salt or c) converting the compd. to an acid addn. salt if the compd. contains a basic N atom. Further, the compds. may be converted to physiol. hydrolyzable esters or to acyloxymethyl or 1-(acyloxy)ethyl esters derived from conventional .beta.-lactam antibiotics. The benzyl ester was prepd. by a) reacting a cyclic anhydride II (q = 0, 2) with HNR1R2 and b) if desired, oxidizing the resulting 6-carbamoyl benzyl ester I (R = CH2Ph, n = 0) to a benzyl ester (n = 1 or 2) with 1 or 2 mol equiv 3-ClC6H4C(O)OOH. II are prepd. by a) reacting 6-dibromo compds. III with 1 mol equiv methylmagnesium Grignard reagent and then with H2C:CHCH2OCOCHO to form allyl ester IV; b) debromination to give V; c) hydrolysis to give the acid VI; and d) reaction with COCl2 in the presence of tertiary amine. Benzyl 6,6-dibromopenicillanate (III, q=0) was treated with MeMgBr at -78.degree., then allyl glyoxalate at -78.degree to give (R)- and (S)-IV (q = 0) the (R)-isomer of which was debrominated to give (S)-V (q = 0). Treating this with BuCHEtCO2Na, then Pd(PPh3)4 gave the Na salt of (S)-VI which was successively treated with COCl2 and NH4OH to give (S)-I (R = which was successively treated with coulz and maken to give (s)-1 (k = CH2Ph, R1 = R2 = H, n = 0). Hydrogenolysis in the presence of NaHCO3 and 10% Pd/C gave (S)-1 (R = Na, R1 = R2 = H, n = 0).

4706-32-5P, N-Glyoxyloylpiperidine 16423-59-9P, N-Glyoxyloylmorpholine 79036-50-3P, N,N-Dimethylglyoxylamide 106435-93-2P, N-Glyoxyloylpyrrolidine 111605-39-1P, N-Isopropylglyoxylamide

RL: SPN (Synthetic preparation); PREP (Preparation)

(prepn. of)

75-16-1, Methylmagnesium bromide TT

RL: RCT (Reactant); RACT (Reactant or reagent) (reaction of, with benzyl dibromopenicillanate and

dimethylglyoxylamide)

16423-59-9, N-Glyoxyloylmorpholine 64370-42-9, Allyl IT 4706-32-5 glyoxalate 79036-50-3, N.N-Dimethylglyoxylamide 106435-93-2, N-Glyoxyloylpyrrolidine 111605-39-1 RL: RCT (Reactant); RACT (Reactant or reagent)

(reaction of, with benzyl dibromopenicillanate and methylmagnesium bromide)

TT 35564-99-9, Benzyl 6,6-dibromopenicillanate RL: RCT (Reactant); RACT (Reactant or reagent) (reaction of, with methylmagnesium bromide and dimethylglyoxylamide)

ANSWER 3 OF 5 CAPLUS COPYRIGHT 2004 ACS on STN L5 1986:552787 CAPLUS AN

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DN

53:1738 OREF 53:227d-f

Glyoxylic acid derivatives

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DN
      105:152787
      Synthesis of psilocin labeled with carbon-14 and tritium
      Poon, Grace; Chui, Yun Cheung; Law, Francis C. P. Dep. Biol. Sci., Simon Fraser Univ., Burnaby, BC, V5A 1S6, Can.
AU
CS
      Journal of Labelled Compounds and Radiopharmaceuticals (1986), 23(2),
      CODEN: JLCRD4; ISSN: 0362-4803
DT
      Journal
      English
LA
      CASREACT 105:152787
               XNMe<sub>2</sub>
                       Ŧ
      14C- and 3H-labeled psilocin (I, X = CH214CH2; C3H2C3H2) tryptamine), the
      principal active agent of hallucinogenic mushrooms, was synthesized from
      2-methyl-3-nitrophenol via 4-benzyloxyindole. 4-Benzyloxygramine was
      treated with K14CN to give 14C-4-benzyloxy-3-indoleacetic acid, an intermediate for I (X = CH214CH2). LiAl3H4 was used to reduce
      4-benzyloxy-3-indole-N,N-dimethylglyoxylamide to give I (X =
      C3H2C3H2).
AB
      14C- and 3H-labeled psilocin (I, X = CH214CH2; C3H2C3H2) tryptamine), the
      principal active agent of hallucinogenic mushrooms, was synthesized from
      2-methyl-3-nitrophenol via 4-benzyloxyindole. 4-Benzyloxygramine was
      treated with K14CN to give 14C-4-benzyloxy-3-indoleacetic acid, an
      intermediate for I (X = CH214CH2). LiAl3H4 was used to reduce
      4-benzyloxy-3-indole-N,N-dimethylglyoxylamide to give I (X =
      C3H2C3II2).
1.5
      ANSWER 4 OF 5 CAPLUS COPYRIGHT 2004 ACS on STN
AN
      1959:28666 CAPLUS
DN
      53:28666
OREF 53:5137e-g
      Glyoxylamide derivatives
ΤI
IN
      Whitfield, Gordon H.
PA
      Imperial Chemical Industries Ltd.
\mathbf{DT}
      Patent
LA
      Unavailable
FAN.CNT 1
      PATENT NO.
                           KIND DATE
                                                     APPLICATION NO. DATE
PΙ
      GB 797604
                                   19580702
                                                     GB
      N,N-Dialkyl substituted glyoxylamide derivs., useful as herbicides were
AB
      prepd. Me2NCOCH(OH)NMe2 (I) (b8 70-3.degree.) (41.2 g.) was dissolved in 50 ml. MeOH and poured into a column (1 1/2'' .times. 3') packed with 500
      g. polystyrenesulfonic acid cation-exchange resin, the resin washed with
      two 500 ml. portions MeOH, and 2 eluate fractions were collected. Removal
      of MeOH from the 1st eluate and distn. of the residue gave 20.69 g. N, N-
      dimethylglyoxylamide Me hemiacetal (II), b20 82.86.degree..
      Similar treatment of the 2nd MeOH eluate gave 3.78 g. II. Exposure of II
      to moist air or treatment with the theoretical amt. of H2O gave
      Me2NCOCHO-0.5H2O (III), m. 121.degree. Similar treatment of I in H2O gave III, m. 121-2.degree., directly. Cf. C.A. 53, 227d.

N,N-Dialkyl substituted glyoxylamide derivs., useful as herbicides were prepd. Me2NCOCH(OH)NMe2 (I) (b8 70-3.degree.) (41.2 g.) was dissolved in 50 ml. MeOH and poured into a column (1 1/2'' .times. 3') packed with 500
      q. polystyrenesulfonic acid cation-exchange resin, the resin washed with
      two 500 ml. portions MeOH, and 2 eluate fractions were collected. Removal
      of MeOH from the 1st eluate and distn. of the residue gave 20.69 g. N,N-dimethylglyoxylamide Me hemiacetal (II), b20 82.86.degree..
      Similar treatment of the 2nd MeOH eluate gave 3.78 g. II. Exposure of 11
      to moist air or treatment with the theoretical amt. of H2O gave
      Me2NCOCHO-0.5H2O (III), m. 121.degree.. Similar treatment of I in H2O gave III, m. 121-2.degree., directly. Cf. C.A. 53, 227d.
      ANSWER 5 OF 5 CAPLUS COPYRIGHT 2004 ACS on STN
L5
      1959:1738 CAPLUS
AN
```

10/620209

=> s e4~e6

1 61960-32-5/BI (61960-32-5/RN) 1 79036-50-3/BI (79036-50-3/RN) 1 939-71-9/BI (939-71-9/RN)

L7

3 (61960-32-5/BI OR 79036-50-3/BI OR 939-71-9/BI)

=> d scan

L7 3 ANSWERS REGISTRY COPYRIGHT 2004 ACS on STN IN 1,3-Benzodioxole-2-carboxamide (6CI, 7CI, 8CI) MF C8 H7 N O3

$$0 \\ C-NH_2$$

\*\*PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT\*\*

HOW MANY MORE ANSWERS DO YOU WISH TO SCAN? (1):2

L7 3 ANSWERS REGISTRY COPYRIGHT 2004 ACS on STN IN Acetamide, 2-hydroxy-2-methoxy-N,N-dimethyl- (9CI) MF C5 H11 N O3

$$\begin{matrix} \text{O} & \text{OH} \\ \parallel & \parallel \\ \text{Me}_{2}\text{N}-\text{C}-\text{CH}-\text{OMe} \end{matrix}$$

\*\*PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT\*\*

L7 3 ANSWERS REGISTRY COPYRIGHT 2004 ACS on STN IN Acetamide, N,N-dimethyl-2-oxo- (9CI) MF C4 H7 N O2

IN Whitfield, Gordon H.

PA Imperial Chemical Industries Ltd.

DT Patent

LA Unavailable

FAN.CNT 1

PATENT NO. KIND DATE

APPLICATION NO. DATE

PI GB 793807

19580423

GB

RIR2NCOCHO, where R1 and R2 are alkyl groups, is prepd. by treating R1R2NCOCHO, where R1 and R2 are alkyl groups, is prepd. by treating R1R2NCO with LiC.tplbond.CH to yield R1R2NCOCH(OLi)NR1R2 followed by hydrolyzing to R1R2NCOCH(OH)NR1R2 (I). I with an acid gives R1R2NCOCHO and the acid salt of R1R2NH. E.g., 219 g. Me2NCO added dropwise to 16 g. LiC.tplbond.CH in 150 ml. boiling MePh, reftuxed 0.5 hr., and MePh and unreacted Me2NCO distd. in vacuo left 108 g. Me2NCOCH(OLi)NMe2 (II). II added to 250 ml. H2O, extd. with ether, dried, and distd. yielded Me2NCOCH(OH)NMe2 (III), b8.0 70-3.degree.. A small portion of III with 2,4-dinitrophenylhydrazine sulfate yielded N,N-dimethylglyoxylamide 2,4-dinitrophenylhydrazone, m. 208.degree.. III is useful as an intermediate in the prepn. of herbicides and pharmaceuticals.

AB RIR2NCOCHO, where R1 and R2 are alkyl groups, is prepd. by treating R1R2NCO with LiC.tplbond.CH to yield R1R2NCOCH(OLi)NR1R2 followed by hydrolyzing to R1R2NCOCH(OH)NR1R2 (I). I with an acid gives R1R2NCOCHO and the acid salt of R1R2NH. E.g., 219 g. Me2NCO added dropwise to 16 g. LiC.tplbond.CH in 150 ml. boiling MePh, reftuxed 0.5 hr., and MePh and unreacted Me2NCO distd. in vacuo left 108 g. Me2NCOCH(OLi)NMe2 (II). II added to 250 ml. H2O, extd. with ether, dried, and distd. yielded Me2NCOCH(OH)NMe2 (III), b8.0 70-3.degree.. A small portion of III with 2,4-dinitrophenylhydrazine sulfate yielded N,N-dimethylglyoxylamide 2,4-dinitrophenylhydrazone, m. 208.degree.. III is useful as an intermediate in the prepn. of herbicides and pharmaceuticals.

10/620209

=> d scan

3 ANSWERS REGISTRY COPYRIGHT 2004 ACS on STN Glycolamide, 2-dimethylamino-N,N-dimethyl- (6CI) C6 H14 N2 O2 IN

MF

CI COM

\*\*PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT\*\*

HOW MANY MORE ANSWERS DO YOU WISH TO SCAN? (1):2

3 ANSWERS REGISTRY COPYRIGHT 2004 ACS on STN Acetic acid, oxo- (9CI)

IN

C2 H2 O3 MF

CI

\*\*PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT\*\*

3 ANSWERS REGISTRY COPYRIGHT 2004 ACS on STN Acetamide, 2-[(2,4-dinitrophenyl)hydrazono]-N,N-dimethyl- (9CI) C10 H11 N5 O5 IN

# 10/620209

## => d his

## (FILE 'HOME' ENTERED AT 17:57:21 ON 16 JAN 2004)

FILE 'CAPLUS' ENTERED AT 18:00:09 ON 16 JAN 2004

E SCHLOEMER G/IN

17 S E4-E6

0 S L1 AND ACETAMIDE?

0 S L1 AND IMIDAZO?

7 S L1 AND PROCESS

Ll

L2 L3

L4

=>

os G1

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ANSWER 1 OF 7 CAPLUS COPYRIGHT 2004 ACS on STN
                  CAPLUS
     2003:222366
     138:238439
     Production of benazepril and analogs by kinetic resolution of an
     intermediate
     Tseng; Wei-Hong; Cheng, Kau-Ming; Schloemer, George; Chen,
     Chien-Wen; Cheng, Chih-Wen
     Scinopharm Taiwan, Ltd., Taiwan U.S. Pat. Appl. Publ., 7 pp., Cont.-in-part of U.S. Ser. No. 910,509.
PΑ
     CODEN: USXXCO
DT
     Patent
     English
LA
FAN.CNT 2
                                              APPLICATION NO.
     PATENT NO.
                        KIND
                              DATE
```

A process for the prepn. of benazepril and analogs I by reaction of compd. II [Z1 = halogen] with compd. III [R1 = H, alkyl or a combination of H and alkyl; R2 = alkyl) in a polar solvent via epimerization and kinetic resoln. of intermediate catalyzed by phase transfer catalyst was developed. Thus, coupling of L-homophenylalanine Et ester to 3-bromo-2,3,4,5-tetrahydro-1H-1-benzapin-2-one using sodium iodide, epimerization and kinetic resoln. of intermediate carboxylic acid, followed by esterification gave compd. (S,S)-I (R1 = H, R2 = Bt) in 80% yield and the ratio of enantiomers detd. by HPLC is SS:RR > 99.5:0.5.

### => d 2-7 bib abs

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ANSWER 2 OF 7 CAPLUS COPYRIGHT 2004 ACS on STN
T.4
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2002:736940 CAPLUS AN

MARPAT 138:238439

DN 137:263201

Process for making taxane derivatives TI

Schloemer, George: Chen, Yung-fa; Lin, Chien Hsin; Daniewski, Wlodzimierz

Scinopharm Taiwan, Ltd., Taiwan PΑ

so U.S. Pat. Appl. Publ., 9 pp.

CODEN: USXXCO

DT Patent English

FAN.	CNT 1 PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
PI	US 2002137955	A1	20020926	US 2001-815517	20010323
	US 6531611	B2	20030311		
	WO 2002076967	A1	20021003	WO 2001-US9348	20010323
	W: CN. JP				
	RW: AT, BE, PT, SE,		, DE, DK, ES,	FI, FR, GB, GR, IE	, IT, LU, MC, NL,

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PRAI US 2001-815517 A 20010323
OS CASREACT 137:263201; MARPAT 137:263201
GI
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\* STRUCTURE DIAGRAM TOO LARGE FOR DISPLAY - AVAILABLE VIA OFFLINE PRINT \* The present invention provides a novel semi-synthetic method of producing a variety of novel taxane derivs. I (R1 = alkoxy, R2 = alkoxy, H; R3 = alkyl; R4 = alkyl, aryl; X = protective group) by reaction of a phenylisoserine deriv. II with a suitably blocked Baccatin III deriv. III. I may be further modified to form pactitaxel and other potentially useful taxane derivs. Thus, III (R1 = R2 = MeO; R3 = Me; R4 = Ph), prepd. from (2R,3S)-phenylisoserine-HCl and .alpha.-methylcinnamic acid, was treated with 7-triethylsilylbaccatin III to give the corresponding I, which was converted to paclitaxel in 4 steps. ANSWER 3 OF 7 CAPLUS COPYRIGHT 2004 ACS on STN 2002:290826 CAPLUS AN 136:310051 DN Process for the preparation of 4,4'-diketo-.beta.-carotene Schloemer, George C.; Schloemer, Danuta A.; Davis, Jeffery L. IN Prodemex, S.A. D.B.C.V., Mex. PA SO U.S., 4 pp. CODEN: USXXAM DT Patent English LΑ FAN.CNT 1 APPLICATION NO. DATE PATENT NO. KIND DATE US 2001-953007 20010913 US 6372946 B1· 20020416 NO 2002-4266 20020906 NO 2002004266 A 20030314 CN 1417207 A 20030514 CN 2002-141620 20020906 A1 20030319 EP 2002-256236 20020909 EP 1293499 R: AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT, IE, SI, LT, LV, FI, RO, MK, CY, AL, TR, BG, CZ, EE, SK 20010913 PRAI US 2001-953007 Α CASREACT 136:310051  $\boldsymbol{\mathsf{A}}$  method of prepg. .beta.-carotene derivs. such as canthaxanthin and astaxanthin was described. The method employs an in situ system to generate hypobromous acid as the oxidizing agent using a salt of sulfite, hydrogen sulfite or bisulfite in combination with a bromate salt. Astaxanthin and canthaxanthin were obtained in good yield with a significantly reduced reaction time. Thus, zeaxanthin was oxidized using sodium hydrogen sulfite in chloroform to form axtaxanthin. THERE ARE 7 CITED REFERENCES AVAILABLE FOR THIS RECORD RE.CNT ALL CITATIONS AVAILABLE IN THE RE FORMAT ANSWER 4 OF 7 CAPLUS COPYRIGHT 2004 ACS on STN AN 2001:798186 CAPLUS DN 135:344616 Oxidative process for the preparation of astaxanthin from zeaxanthin using a halogenating agent with the salt of chloric or bromic acid in an inert solvent IN Schloemer, George C.; Davis, Jeffery L. Prodemex, S.A. de C.V., Mex. SO PCT Int. Appl., 12 pp. CODEN: PIXXD2 DT Patent LA English FAN.CNT 1 KIND DATE APPLICATION NO. DATE PATENT NO. WO 2001081301 A2 20011101 WO 2001-US13295 20010425 AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, BZ, CA, CH, CN, CR, CU, CZ, CZ, DE, DE, DK, DK, DM, DZ, EE, EE, ES, FI, FI, GB, GD, GE, GH, GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT, LU, LV, MA, MD, MG, MK, MN, MW, MX, MZ, NO,

NZ. PL, PT, RO, RU, SD, SE, SG, SI, SK, SK, SL, TJ, TM, TR, TT, TZ, UA, UG, US, UZ, VN, YU, ZA, ZW, AM, AZ, BY, KG, KZ, MD, RU,

US 2001-813685

20010319

RW: GH, GM, KE, LS, MW, MZ, SD, SL, SZ, TZ, UG, ZW, AT, BE, CH, CY, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, TR, BF, BJ, CF, CG, CI, CM, GA, GN, GW, ML, MR, NE, SN, TD, TG

TJ. TM

**A**1

20011213

US 2001051357

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US 6376717
                       B2
                            20020423
                                           EP 2001-932633
                                                             20010425
                            20030122
     BP 1276719
                       A2
        R: AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT, IE, SI, LT, LV, FI, RO, MK, CY, AL, TR
                                            NO 2001-6293
                                                             20011220
     NO 2001006293
                      Α
                            20020211
                                                             20011221
     ZA 2001010503
                            20020829
                                            ZA 2001-10503
PRAI US 2000-199875P
                       P
                            20000426
     US 2001-813685
                       Α
                            20010319
    WO 2001-US13295
                       W
                            20010425
    CASREACT 135:344616
OS
    Astaxanthin is prepd. from zeaxanthin by oxidn. using a halogenating agent
AB
     with the salt of chloric or bromic acid in an inert solvent.
     ANSWER 5 OF 7 CAPLUS COPYRIGHT 2004 ACS on STN
T.4
     1996:689881 CAPLUS
AN
DN
     126:19174
     Preparation of acyclovir using 1,3 dioxolane
ΤI
     Schloemer, George C.; Han, Yeun-kwei; Harrington, Peter J.
IN
     Syntex (U.S.A.) Inc., USA
PA
     U.S., 8 pp., Cont.-in-part of U.S. Ser. No. 280,269, abandoned.
SO
     CODEN: USXXAM
DT
     Patent
     English
LA
FAN.CNT 2
     PATENT NO.
                      KIND DATE
                                            APPLICATION NO. DATE
                                            US 1995-426005
                                                             19950427
PΙ
     US 5567816
                       Α
                            19961022
                                            CA 1995-2152863 19950628
     CA 2152863
                       AA
                            19960127
                                            JP 1995-176022
     JP 08053451
                       A2
                            19960227
                                                             19950712
     EP 709385
                       A1
                            19960501
                                            EP 1995-110955
                                                             19950713
        R: AT, BE, CH, DE, DK, ES, FR, GB, GR, IE, IT, LI, LU, NL, PT, SE
                                            CN 1995-115316
     CN 1122805
                       A
                            19960522
                                                             19950725
     BR 9503442
                       A
                            19960604
                                            BR 1995-3442
                                                             19950725
     FI 9503580
                            19960127
                                            FI 1995-3580
                                                             19950726
PRAI US 1994-280269
                            19940726
     US 1995-426005
                            19950427
    A process for the prepn. of acyclovir via coupling of guanine or
     silylated guanines with 1,3-dioxolane in the presence of a selective
     alkylation catalyst selected from the group consisting of
     trifluoromethanesulfonic acid, trimethylsilyl trifluoromethanesulfonate,
     and bistrimethylsilyl sulfonate, and hydrolyzing the product thus formed.
     ANSWER 6 OF 7 CAPLUS COPYRIGHT 2004 ACS on STN
L4
     1996:313501 CAPLUS
AN
DN
     124:343989
     Method for producing 9-(2-hydroxyethoxymethyl)guanine (acyclovir) as
TI
     antiviral agent
     Han, Yuen-Kwei; Harrington, Peter John; Schloemer, George Charles
IN
     F. Hoffmann-La Roche Ag, Switz.
PA
     Can. Pat. Appl., 28 pp.
so
     CODEN: CPXXEB
חד
     Patent
     English
LA
FAN.CNT 2
                                            APPLICATION NO.
                                                             DATE
     PATENT NO.
                      KIND
                            DATE
                            19960127
                                            CA 1995-2152863
                                                             19950628
     CA 2152863
                       AA
PI
                                            US 1995-426005
                                                             19950427
                            19961022
     US 5567816
                       Α
PRAI US 1994-280269
                            19940726
     US 1995-426005
                            19950427
     CASREACT 124:343989; MARPAT 124:343989
OS
GI
```

AB An efficient and selective process for the synthesis of the antiviral 9-(2-hydroxyethoxymethyl) guanine (acyclovir) (I) involves (1) contacting a silylated guanine or mixts. of silylated guanine (II; Z1, Z2, Z3 = H, R1R2R3Si; wherein R1 - R3 = lower alkyl; provided that at least one of Z1 - Z3 = R1R2R3Si) with 1,3-dioxolane (III) in the presence of a selective alkylation catalyst and (2) hydrolyzing the product formed. Said catalyst is selected form CF3SO3H, CF3SO3SiMe3, and bis(trimethylsilyl) sulfonate and CF3SO3HMe3; is generated by contacting CF3SO3H with hexamethyldisilazane. The process avoids the use of acyl groups for protection of guanine, essentially specific for the prepn. of the N-9 isomer, thus eliminates the need for the chromatog, sepn. of the N-9/N-7 isomer mixts., provides I in good yields, requires simple starting materials and reaction conditions, and is carried out from start to finish in a single reaction vessel. Thus, a mixt. of 25 g guanine, 125 mL hexamethyldisilazane, and 1 mL CF3SO3SiMe3 was refluxed at 130-135 degree. for 24 h, cooled to 70 degree., treated with 25 mL 1,3-dioxolane, refluxed for 16 h, distd. under reduced pressure to remove excess hexamethyldisilazane, cooled to 70 degree., poured into a mixt. of 600 mL 10% aq. AcOH, and heated to give a soln. The soln. was treated with 1.25 g activated carbon to remove any color, filtered, and the filtrate was slowly cooled to 5 degree. to give, after filtering off the white cryst. solid formed, 78% I.

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L4 ANSWER 7 OF 7 CAPLUS COPYRIGHT 2004 ACS on STN
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AN 1989:23725 CAPLUS

DN 110:23725

TI Process for preparing (.+-.)-1,2-dihydro-3H-pyrrolo[1,2-a]pyrrole-1,7-dicarboxylates as intermediates for pharmaceuticals

IN Khatri, Hiralal N.; Fleming, Michael P.; Schloemer, George C.

PA Syntex (U.S.A.), Inc., USA SO Eur. Pat. Appl., 28 pp.

SO Eur. Pat. Appl., 28 pp CODBN: EPXXDW

DT Patent

LA English

FAN.CNT 1

FAN.					
					APPLICATION NO. DATE
ΡI					EP 1988-100390 19880113
	ΕP	275092	B1	19920603	
					GB, GR, IT, LI, LU, NL, SE
					US 1987-3162 19870114
	US	4849526	Α.	19890718	US 1987-3104 19870114
	DK	8800143	A	19880715	DK 1988-143 19880113
	FI	8800133		19880715	FI 1988-133 19880113
		90344		19931015	
	FΙ	90344 -		19940125	
	NO	8800127	A	19880715	NO 1988-127 19880113
	NO	169124	В	19920203	
		169124		19920513	
	ΑU	8810240	A1	19880721	AU 1988-10240 19880113
	ΑU	613334	B2	19910801	
	JР	63198684			JP 1988-6757 19880113
		8800225			ZA 1988-225 19880113
	HU	48881	A2	19891128	HU 1988-117 19880113
		200606		19900728	
	HU	51595	A2	19900528	HU 1989-5354 19880113
	HU	201728	В	19901228	l e e e e e e e e e e e e e e e e e e e
	HU	52045	A2	19900628	HU 1989-5355 19880113
	HU	203532	В	19910828	
	ΗU	52046	A2	19900628	HU 1989-5356 19880113

	HƯ	203721	В	19910930			,
	IL	85094	A1	19910916	IL	1988-85094	19880113
	IL	96388	A1	19910916	IL	1988-96388	19880113
	ΙL	96389	A1	19910916	IL	1988-96389	19880113
	ΑT	76873	E	19920615	ΤA	1988-100390	19880113
	ES	2041703	<b>T</b> 3	19931201	ES	1988-100390	19880113
	HU	213614	В	19970828	HŲ	1990-1266	19880113
	CA	1340404	A1	19990223	CA	1988-556465	19880113
	US	4874872	A	19891017	US	1988~255799	19881011
	US	4937368	A	19900626	US	1989-299701	19890123
	NO	9003993	A	19880715	NO	1990-3993	19900913
	NO	174583	В	19940221			
	NO	174583	С	19940601			
	NO	9003994	A	19880715	NO	1990-3994	19900913
	NO	174346	В	19940110			
	NO	174346	С	19940420			
	NO	9003995	A	19880715	NO	1990-3995	19900913
	NO	173828	В	19931101			
	NO	173828	C	19940209			_
	FI	92488	В	19940815	FΙ	1991-2709	19910605
	FI	92488	C	19941125			
	FI	95242	В	19950929	ΡI	1991-2710	19910605
	FI	95242	С	19960110			
	FΙ	91148	В	19940215	FI	1993-320	19930126
	FI	91148	C	19940525			
PRAI	US	1987-3104	A	19870114			
	US	1987-3162	A	19870114			
	EΡ	1988-100390	A	19880113			
	HU	1988-117	A	19880113			
		1988-85094	A	19880113			
		1988-127	Al	19880113			
os	CA:	SREACT 110:2372	5; MA	RPAT 110:23725			
GT							

AB A process for producing diesters I (R = alkyl), useful as intermediates for pharmaceuticals II [Ar = alkyl, alkoxy, or halo (un)substituted Ph, 2- or 3- furoyl, 2- or 3-thienyl, 2- or 3-pyrryl; R = H, alkyl] useful as analgesics, antiinflammatories, antipyretics, and smooth muscle relaxants (no data), comprised: a) cyclizing pyrrole III (X = halo) with a hindered amine in an aprotic polar solvent; or b) reacting pyrrolidine IV with XCH2CHo (X = halo) in aq. soln. I (R = alkyl) are sapond. to I (R = H) which are monoesterified to I (R at 1 = alkyl, R at 7 = H) which are decarboxylated to II (no ArCO group). These are aroylated with an amide or morpholide to give II. 1 (R = H), which had been prepd. in 5 steps from BrCH2CH2NH2. HBr and (MeO2CCH2)2CO was converted in 4 steps into II (Ar = 4-MeC6H4, R = H).

L Number	Hits	Search Text	DB	Time stamp
1	3478	phosphorus adj tribromide	USPAT;	2004/01/16 17:19
			US-PGPUB	
2	30687	thionyl adj chloride	USPAT;	2004/01/16 17:20
			US-PGPUB	Ì
3	315	(phosphorus adj tribromide) near (thionyl adj chloride)	USPAT;	2004/01/16 17:20
			US-PGPUB	
4	593	imidazopyridine .	USPAT;	2004/01/16 17:21
			US-PGPUB	
5	0	((phosphorus adj tribromide) near (thionyl adj chloride)) and	USPAT;	2004/01/16 17:20
		imidazopyridine	US-PGPUB	
6	271	imidazopyridines	USPAT;	2004/01/16 17:21
			US-PGPUB	
7	745	imidazopyridine or imidazopyridines	USPAT;	2004/01/16 17:21
		·	US-PGPUB	_
8	0	(imidazopyridine or imidazopyridines) and ((phosphorus adj	USPAT;	2004/01/16 17:22
		tribromide) near (thionyl adj chloride))	US-PGPUB	
9	11485	halogenating	USPAT;	2004/01/16 17:22
			US-PGPUB	
10	0	((phosphorus adj tribromide) near (thionyl adj chloride)) near	USPAT;	2004/01/16 17:22
		halogenating	US-PGPUB	
11	180	((phosphorus adj tribromide) near (thionyl adj chloride)) same	USPAT;	2004/01/16 17:45
		halogenating	US-PGPUB	
12	3	(((phosphorus adj tribromide) near (thionyl adj chloride)) same	USPAT;	2004/01/16 17:47
		halogenating) and sleep	US-PGPUB	
13	113	hydrolysis and (((phosphorus adj tribromide) near (thionyl adj	USPAT;	2004/01/16 17:47
		chloride)) same halogenating)	US-PGPUB	0004/04/40 47 47
14	310	hydrolysis same (phosphorus adj tribromide) (((phosphorus adj	USPAT;	2004/01/16 17:47
		tribromide) near (thionyl adj chloride)) same halogenating)	US-PGPUB	0004/04/40 477 40
15	1	hydrolysis same (((phosphorus adj tribromide) near (thionyl adj	USPAT;	2004/01/16 17:48
'	1	chloride)) same halogenating)	US-PGPUB	<u> </u>